

due at this time, but applicants authorize the Commissioner to charge any fees now due to Arnold,
White & Durkee Deposit Acct. No. 01-2508/CADL:002/PAR.

AMENDMENTS

In the Claims:

Please amend the claims as follows:

Sub 13

63. (Amended) [An] The antigen composition according to claim 62, wherein UTAA is purified at least about 100-fold over UTAA found in urine.

64. (Amended) [An] The antigen composition according to claim 62, wherein said UTAA is present as at least about 0.6% of total protein in said composition.

65. (Amended) [A] The method of claim 19, [for] wherein said method comprises enhancing in a subject the production of antibodies reactive with UTAA [comprising administering an effective amount of the antigen composition of claim 62].

Please add the following claims:

25 Sub 4

66. The composition of claim 63, wherein said UTAA is purified 105-fold over UTAA found in urine.

Sub 5

67. The composition of claim 62, wherein said UTAA has an isoelectric point of about 6.1.

68. The composition of claim 62, wherein said UTAA is heat stable at 100°C.

69. The composition of claim 62, wherein said UTAA is about 95% free of immunoglobulin.

Sub 5

70. The composition of claim 62, wherein said UTAA is about 99.5% free of immunoglobulin.

Sub 2

71. The composition of claim 62, wherein said UTAA contains glycosidase-sensitive carbohydrates.

72. The method of claim 65, wherein the observed enhancement of antibody production is about 2- to 5-fold.

Sub 6

73. A pharmaceutical composition comprising (i) an antigen composition comprising a substantially purified tumor antigen, wherein the tumor antigen is identified as comprising Urinary Tumor Associated Antigen (UTAA) subunit which, after reduction by β -mercaptoethanol and separation by SDS-polyacrylamide gel electrophoresis, exhibits a molecular weight of about 90 to 100 kD and (ii) a pharmaceutical buffer.

74. The pharmaceutical composition of claim 74, wherein said antigen composition is present as at least about 0.63 μ g/ml of buffer.

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75. The pharmaceutical composition of claim 75, wherein said antigen composition is present as at least about 1.4 $\mu\text{g}/\text{ml}$ of buffer.

76. The pharmaceutical composition of claim 76, wherein said antigen composition is present as at least about 36 $\mu\text{g}/\text{ml}$ of buffer.

77. The pharmaceutical composition of claim 75, wherein said antigen composition is present as at least about 40 $\mu\text{g}/\text{ml}$ of buffer.

78. The pharmaceutical composition of claim 75, wherein said antigen composition is present as at least about 100 $\mu\text{g}/\text{ml}$ of buffer.

79. The pharmaceutical composition of claim 75, wherein said antigen composition is present as at least about 200 $\mu\text{g}/\text{ml}$ of buffer.
